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APPLICATION NUMBER	FILING DATE		FIRST NAMED APPLICANT		ATTORN	EY DOCKET NO.
09/026,276	02/19/98	KENTEN			J	16N-9601
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				DATE MAIL		<b>)</b> 12/30/98
		•				12/30/50
This is a communication from COMMISSIONER OF PATE	n the examiner in charg NTS AND TRADEMARI	e of your application	on. - ,			
	. 01	FICE ACTIO	N SUMMARY			
Responsive to communicat	ion(s) filed on	2-19-	38			
This action is FINAL.						
accordance with the practic shortened statutory period for hischever is longer, from the n le application to become abai .136(a).	or response to this ac	tion is set to exp	oire	vithin the per	iod for re	, or thirty days, seponse will cause rovisions of 37 CFR
Isposition of Claims						
Claim(s)	1-100				is/are p	ending in the application
Of the above, claim(s)						
☐ Claim(s)						
Claim(s)						
Claim(s)						is/are objected to.
☐ Claim(s)	- 100		ar	e subject to	restrictio	on or election requiremen
Application Papers						
☐ See the attached Notice	of Draftsperson's Pa	atent Drawing R	eview, PTO-948.			
☐ The drawing(s) filed on						
☐ The proposed drawing of	correction, filed on			i	s 🗌 ap	proved  disapprove
☐ The specification is obje	ected to by the Exam	iner.				
☐ The oath or declaration	is objected to by the	Examiner.				
Priority under 35 U.S.C. §	119					
Acknowledgement is mad	e of a claim for foreig	n priority under	35 U.S.C. § 119(a)	)-(d).		
☐ All ☐ Some* ☐ N	one of the CERTIF	TED copies of t	ne priority document	s have been	I	
received.						
received in Application	on No. (Series Code/	Serial Number)			<u>-</u> ·	
received in this nation	nal stage application	from the Interna	itional Bureau (PCT	Rule 17.2(a	i)). ·	
*Certified copies not receiv						•
Acknowledgement is made	le of a claim for dom	estic priority und	ler 35 U.S.C. § 119	)(e).		
Attachment(s)						
☐ Notice of Reference Ci	ted, PTO-892					
Information Disclosure	Statement(s), PTO-1	449, Paper No(	s)			
☐ Interview Summary PT	ΓΟ-413					

☐ Notice of Draftsperson's Patent Drawing Review, PTO-948

PTOL-326 (Rev. 10/95)

Notice of Informal Patent Application, PTO-152 Sequence rules.

Notice to Complete See OFFICE ACTION ON THE FOLLOWING PAGES -

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## **DETAILED ACTION**

## Election/Restriction

- 1. Restriction to one of the following inventions is required under 35 U.S.C. 121:
  - I. Claims 1-75, 81-83 drawn to a DNA construct encoding a ubiquitin fusion protein and the protein encoded thereby, classified in class 530, subclass 350.
  - II. Claims 76-79, 84-85, 87-93 drawn to a method for stimulating an immune response in an animal, the immune response being directed to a ubiquitin fusion protein or reducing levels of a predetermined protein in an animal, by providing a ubiquitin fusion protein, classified in class 514, subclass 2.
  - III. Claims 80, 94-100 drawn to a method for stimulating an immune response in an animal, the immune response being directed to a ubiquitin fusion protein or reducing levels of predetermined protein in an animal, by providing DNA construct encoding a ubiquitin fusion protein, classified in class 514, subclass 44.
  - IV. Claim 86, drawn to a method for identification of antibodies in experimental or diagnostic samples by incubating a ubiquitin fusion protein with antibodies, and detecting the binding of these antibodies to ubiquitin fusion protein, classified in class 435, subclass 7.1.

The inventions are distinct, each from the other because of the following reasons:

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Inventions I and II are related as product and process of use. The inventions can be shown to be distinct if either or both of the following can be shown: (1) the process for using the product as claimed can be practiced with another materially different product or (2) the product as claimed can be used in a materially different process of using that product (M.P.E.P. § 806.05(h)). In the instant case the ubiquitin fusion protein can be used diagnostically (e.g in screeing).

Inventions I and III are related as product and process of use. The inventions can be shown to be distinct if either or both of the following can be shown: (1) the process for using the product as claimed can be practiced with another materially different product or (2) the product as claimed can be used in a materially different process of using that product (M.P.E.P. § 806.05(h)). In the instant case the DNA as claimed can be used as a hybridization probe.

Inventions II-IV are independent and distinct, each from the other, because the methods are practiced with materially different process steps for materially different purposes and each method requires a non-coextensive search because of different starting materials, process steps and goals.

Having shown that these inventions are distinct for the reasons given above and have acquired a separate status in the art as shown by their different classification and recognized divergent subject matter as defined by MPEP § 808.02, the Examiner has prima facie shown a serious burden of search (see MPEP § 803). Therefore, an initial requirement of restriction for examination purposes as indicated is proper.

2. This application contains claims directed to the following patentably distinct species of the claimed invention: a ubiquitin fusion protein fused to one of the following epitopes:

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(a) gonadotropin releasing hormone. (b) tumor necrosis factor. (c) immunoglubulin. (d) human immunodeficiency virus proteins. (e) chorionic gonadotropin. (f) inhibin. (g) growth hormones and (h) sperm proteins.

Applicant is required under 35 U.S.C. 121 to elect a single disclosed species for prosecution on the merits to which the claims shall be restricted if no generic claim is finally held to be allowable. Currently, claims 1-14, 16-34, 36-54, 56-71, 73-89, 93-96, 100 are generic.

Applicant is advised that a reply to this requirement must include an identification of the species that is elected consonant with this requirement, and a listing of all claims readable thereon, including any claims subsequently added. An argument that a claim is allowable or that all claims are generic is considered nonresponsive unless accompanied by an election.

Upon the allowance of a generic claim, applicant will be entitled to consideration of claims to additional species which are written in dependent form or otherwise include all the limitations of an allowed generic claim as provided by 37 CFR 1.141. If claims are added after the election, applicant must indicate which are readable upon the elected species. MPEP § 809.02(a).

Should applicant traverse on the ground that the species are not patentably distinct, applicant should submit evidence or identify such evidence now of record showing the species to be obvious variants or clearly admit on the record that this is the case. In either instance, if the examiner finds one of the inventions unpatentable over the prior art, the evidence or admission may be used in a rejection under 35 U.S.C. 103(a) of the other invention.

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3. Applicant is advised that the response to this requirement to be complete must include an election of the invention to be examined even though the requirement be traversed (37 CFR 1.143).

Applicant is reminded that upon the cancellation of claims to a non-elected invention, the inventorship must be amended in compliance with 37 C.F.R. § 1.48(b) if one or more of the currently named inventors is no longer an inventor of at least one claim remaining in the application. Any amendment of inventorship must be accompanied by a diligently-filed petition under 37 C.F.R. § 1.48(b) and by the fee required under 37 C.F.R. § 1.17(h).

This application contains sequence disclosures that are encompassed by the definitions for nucleotide and/or amino acid sequences set forth in 37 CFR 1.821(a)(1) and (a)(2). However, this application fails to comply with the requirements of 37 CFR 1.821 through 1.825 for the reason(s) set forth on the attached Notice To Comply With Requirements For Patent Applications Containing Nucleotide Sequence And/Or Amino Acid Sequence Disclosures.

Applicant is given ONE MONTH, or THIRTY DAYS, whichever is longer, from the mailing date of this letter within which to comply with the sequence rules, 37 CFR 1.821 - 1.825. Failure to comply with these requirements will result in ABANDONMENT of the application under 37 CFR 1.821(g). Extensions of time may be obtained by filing a petition accompanied by the extension fee under the provisions of 37 CFR 1.136(a). In no case may an applicant extend the period for reply beyond the SIX MONTH statutory period. Direct the reply to the undersigned. Applicant is requested to return a copy of the attached Notice to Comply with the reply.

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**Advisory Information** 

Any inquiry concerning this communication or earlier communications from the examiner

should be directed to Fozia Hamud whose telephone number is (703) 308-8896. The examiner can

normally be reached on Monday-Friday from 8:00AM to 4:30PM (Eastern time).

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor,

Lila Feisee, can be reached on (703) 308-2731.

Official papers filed by fax should be directed to (703) 308-4227. Faxed draft or informal

communications with the examiner should be directed to (703) 308-0294.

Any inquiry of a general nature or relating to the status of this application or proceeding should be

directed to the Group receptionist whose telephone number is (703) 308-0196.

Fozia Hamud FH

Patent Examiner

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December 29, 1998

Application No.

Please return a copy of this notice with your response.

For PatentIn software help, call (703) 557-0400